

REMARKS

The title has been amended to conform with the English translation of International Patent Application No. PCT/JP2005/002994.

The specification has been amended in accordance with 37 CFR §1.78 to make reference to the International Application from which this application originates and to incorporate by reference the Japanese priority application. The specification has also been amended to correct informalities and typographical errors. A mark up copy and a clean copy of the substitute specification are provided. No new matter has been added.

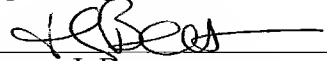
Claims have been amended to correct informalities and typographical errors, and to eliminate multiple dependency. The amendment is made to reduce filing fees and not for any other reason related to patentability of such claims. No new matter has been added.

The abstract has been amended to conform to U.S. practice. No new matter has been added.

The claim fee was calculated based on the amended claims above. Please examine the application in view of the amendments set forth above.

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Respectfully submitted,

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SPECIFICATION

5 METHOD FOR FORMING POWDER MOLDING PRODUCT AND MOLD APPARA-
TUS FOR POWDER MOLDING

Cross Reference to Prior Application

This is a U.S. national phase application under 35
10 U.S.C. §371 of International Patent Application No.
PCT/JP2005/002994 filed February 24, 2005 and claims the
benefit of Japanese Application No. 2004-055363, filed Feb-
ruary 27, 2004. The International Application was pub-
lished in Japanese on September 9, 2005 as International
15 Publication No. WO 2005/082562 A1 under PCT Article 21(2).

Field of the Invention

The present invention relates to a method for forming
a powder molding product by filling raw powders in a mold
20 for powder molding, and also relates to a mold apparatus
for such powder molding.

Background Art

A compact powder, which is used for the production of
25 sintered products, is formed by pressing raw powders such

as Fe-based powders, Cu-based powders or the like in a mold, and then a sintered body is formed through a sintering process. And in the molding process, the molding product undergoes a press-molding process, using a mold. At the
5 time of the press-molding, however, a friction between a molding product and a mold is generated. For this reason, when mixing powders, a water insoluble fatty acid lubricant, such as zinc stearate, calcium stearate, lithium stearate, etc., is added so as to impart lubricity.

10 However, the method of applying a lubricant to raw powders has limitations of improvement of the density of a molding product. Accordingly, in order to obtain a high-density molding product, there is proposed a method for forming a powder molding product which can make up for the
15 lack of lubricity by applying the same lubricant as the one added to raw powders to a mold while reducing the amount of lubricant added to raw powders (~~for~~for example, see Patent Document 1).

This conventional method of molding is a method for
20 forming powder molding product which comprises steps of applying water dispersed in a high fatty acid lubricant to an inner surface of a heated mold and press-molding metal powders by filling the metal powders in the mold and pressing the same at such a pressure that the high fatty acid lubri-
25 cant is chemically bonded to the metal powders so as to

produce a film of metallic soap, wherein the mold is heated,
and the inner surface thereof is coated with the high fatty
acid lubricant such as lithium stearate; heated metal pow-
ders are filled into this mold and are subjected to press-
5 molding at such pressure that the high fatty acid lubricant
is chemically bonded to the metal powders so as to produce
the film of metallic soap, whereby the film of metallic
soap is produced on the inner surface of the mold to
thereby reduce the friction between the molding product of
10 the metallic powders and the mold, thereby enabling the re-
duction of pressure for ejecting the molding product.

As the fact that the same lubricant as one added to
the raw powders is used for the mold results in the use of
the water insoluble lubricant, the lubricant applied to the
15 mold is applied in a solid powder state. For this reason,
other lubricant application methods are also known, such as
electrostatic application of lubricant powders or dry ap-
plication of lubricant which is dispersed in water by sur-
factant.

20 ~~Patent Document 1: Japanese Registered Patent Publi-~~
~~cation No. 3303970 (see paragraphs 0012 and 0013).~~

DisclosureSummary of the invention

25 ~~Problems to be solved by the invention~~

According to the conventional art disclosed in the
~~above Patent Document 1, Japanese Registered Patent Publi-~~
~~cation No.3309970~~ since the lubricant dispersed in water is
applied to the mold in a state of solid powders, that is,
5 in such state that the solid powders of the lubricant are
dispersed and mixed in water, a fine and uniform film can-
not be formed, and thus there is a problem that producing a
molding product of a stable quality is difficult.

The present invention has been made to solve the
10 above problems. The same applicant proposes in the Japa-
nese Patent Application No. 2002-338621, a method for form-
ing a powder molding product by filling a molding portion
formed in a mold body with a raw powder and then fitting
punches into the molding portion, comprising the steps of
15 applying an aqueous solution obtained by dissolving a lu-
bricant in solvent to the molding portion prior to filling
the molding portion with a raw powder, and evaporating the
aqueous solution thus applied to thereby form a crystal-
lized layer on the molding portion which enables producing
20 a fine and uniform layer of lubricant on a molding portion
by said crystallized layer. And optimum aqueous solution
is obtained by the more and more development of crystal-
lized layer.

It is, accordingly, an object of the present inven-
25 tion to provide a method for forming a powder molding prod-

uct and mold apparatus for powder molding which enables the stable production of a high density powder molding product by forming a fine and uniform film of lubricant on a molding portion.

5

~~Problems to be Solved by the Invention~~

~~Claim 1~~An aspect of the present invention is a method for forming a powder molding product by filling a molding portion formed in a mold body with a raw powder and then fitting punches into the molding portion, which is characterized in that applying an aqueous solution obtained by dissolving a water soluble lubricant having at least 3g of solubility for 100 g of water at 20°C in a water to the molding portion prior to filling the molding portion with a raw powder, and evaporating the aqueous solution thus applied to thereby form a crystallized layer on the molding portion.

Further, ~~the present invention of claims 2-12 is embodied~~ embodiments are that said lubricants are used at least one selected from a group of oxo-acid based metal salts or organic acid based metal salts. ~~Said~~In an embodiment aqueous solution ~~of claim 13 of the present invention is~~ completely dissolved in water to have a concentration greater than or equal to 0.01 % by weight concentration but less than saturated concentration. ~~And and the present invention of~~

~~claim 14 is said lubricant is potassium salt or sodium salt.~~

Further, ~~the present invention of claim 15 is that~~
~~embodiments add an antiseptic substance, is added into said~~
~~lubricant. The present invention of claim 16 is that~~ a de-
5 foaming agent ~~is added into said lubricant. Furthermore,~~
~~the present invention of claim 17 is that~~ and a water solu-
ble solvent ~~is added into said lubricant. Also the present~~
~~invention of claim 18 is that said aqueous solvent is~~ can be
alcohol or ketone. Further, ~~the present invention of claim~~
10 ~~19 is that no halogen elements is included~~ can be excluded
~~from is the lubricant.~~

With regard to a method for forming a powder molding
product, for example, by completely dissolving water solu-
ble phosphate based metal salt such as dipotassium hydrogen
15 phosphate, disodium hydrogen phosphate in water to have a
concentration greater than or equal to 0.01 % by weight but
less than a saturated ~~temperature~~ concentration at a fric-
tion surface of molding product, to become a uniform phase,
then applying a solution to a surface of a molding portion,
20 and evaporating, the crystal of the lubricant grows to form
a crystallized layer.

Further, an embodiment of the invention of claim 20
is a mold apparatus for powder molding, ~~comprising~~ having a
mold body with a through-hole for forming a side of a pow-
25 der molding product, a lower punch to be fitted into the

through-hole from beneath, an upper punch to be fitted into the through-hole from above, a spray member from which a lubricant aqueous solution is faced into the through-hole, a heater provided around a molding portion of the powder molding product, the molding portion being defined by the through-hole and the lower punch which is fitted into the through-hole, and a temperature control system keeping a temperature of the heater higher than an evaporating temperature of the aqueous solution.

10 Furthermore, ~~the invention of claim 21~~ another embodiment is a mold apparatus for powder molding, ~~comprising~~ having a mold body with a through-hole for molding a side of a powder molding product, a lower punch to be fitted into the through-hole from beneath, an upper punch to be fitted into the through-hole from above, a spray member from which a lubricant aqueous solution is faced into the through-hole, a heater provided around a molding portion of the powder molding product, the molding portion being defined by the through-hole and the lower punch which is fitted into the through-hole, a temperature control system keeping a temperature of the heater higher than an evaporating temperature of the aqueous solution, but lower than a melting temperature of said lubricant.

25 According to the ~~constitution of claim 20 or 21~~ above, an aqueous solution in which water soluble lubricant having

at least 3g of solubility for 100 g of water at 20°C is applied to the heated molding portion prior to a raw powder being filled in the molding portion defined by the through-hole in the mold body and the lower punch to be fitted into the through-hole, so that the solution is evaporated to thereby form a fine crystallized layer around the surface of the molding portion. Thereafter, the molding portion is filled with a raw powder, and then the upper punch is fitted from above into the through-hole, to thereby form a powder molding product.

~~Effect of the Invention~~

~~Claim 1 of the~~ The present invention ~~is~~ can be a method for forming a powder molding product by filling a molding portion formed in a mold body with a raw powder and then fitting punches into the molding portion, which is characterized in that applying an aqueous solution obtained by dissolving a lubricant in solvent to the molding portion prior to filling the molding portion with a raw powder, and evaporating the aqueous solution thus applied to thereby form a crystallized layer on the molding portion. By forming more fine and uniform crystallized layer for lubrication, the reduction of a pressure for ejecting the powder molding product as well as the improvement of the density of the powder molding product can be obtained.

Further, ~~claim 2~~an aspect of the invention is that the lubricant is oxo-acid based metal salt.

Further, ~~claim 3~~an embodiment of the present invention is that said lubricant is phosphate metal salt, sulfate metal salt, borate metal salt, silicate metal salt, tungstate metal salt, organic acid based metal salt, nitrate metal salt or carbonate metal salt. ~~Claim 4 of the present invention is that~~Alternatively, the said lubricant is dipotassium hydrogen phosphate, disodium hydrogen phosphate, tripotassium phosphate, trisodium phosphate, potassium polyphosphate, sodium polyphosphate, riboflavin potassium phosphate and riboflavin sodium phosphate. ~~Claim 5 of the present invention is that said~~Also, the lubricant ~~can~~be potassium sulfate, sodium sulfate, potassium sulfite, sodium sulfite, potassium thiosulfate, sodium thiosulfate, potassium dodecyl sulfate, sodium dodecyl sulfate, potassium dodecyl benzen sulfate, sodium dodecyl benzene sulfate, Food Blue No.1., Food Yellow No.5., potassium ascorbyl sulfate ester and sodium ascorbyl sulfate ester.

~~Claim 6 of the present invention~~An embodiment is that ~~said~~the lubricant is potassium tetraborate or sodium tetraborate. ~~Claim 7 of the present invention is that said~~Alternately, the lubricant is potassium silicate, ~~or~~ sodium silicate. ~~Claim 8 of the present invention is that said lubricant is~~ potassium tungstate, ~~or~~ sodium tungstate.

~~Claim 9 of the present invention is that said lubricant is,~~
potassium acetate, sodium acetate, potassium benzoate, so-
dium benzoate, dipotassium terephthalate, disodium
terephthalate, potassium ascorbate, or sodium ascorbate.

5 ~~Claim 10~~ Also, an embodiment of the present invention is
that ~~said~~ the lubricant is potassium nitrate, ~~or~~ sodium ni-
trate, ~~Claim 11 of the present invention is that said~~
~~lubricant is~~ potassium carbonate, sodium carbonate, potas-
sium hydrogen carbonate, or sodium hydrogen carbonate.

10 ~~Claim 12 of the present invention is that said~~ Furthermore,
the lubricant uses one or more lubricants described ~~in~~
~~claims 2 to 11 above.~~ ~~Claim 13 of the present invention is~~
~~that said~~ The aqueous solution ~~is~~ can be ~~the~~ one in which
said lubricant is completely dissolved in water to have a
15 concentration greater than or equal to 0.01 % by weight but
less than saturated concentration. ~~Claim 14 of the present~~
~~invention is that the~~ The lubricant ~~is~~ can be potassium salt
or sodium salt. ~~Claim 15 of the present invention is that~~
~~and an antiseptic substance is added into said lubricant.~~

20 ~~Claim 16 of the present invention is that~~ or a defoaming
agent ~~is~~ can be added into the lubricant. Further, ~~claim 17~~
~~of the present invention is that a~~ water soluble solvent is
added into the lubricant. ~~Claim 18 of the present invention~~
~~is that and the~~ said solvent is alcohol or ketone. ~~Claim 19~~
25 ~~of the present invention is that~~ Also, an embodiment is that

no halogen element is included in the lubricant.

~~Further claim 20 and claim 21 of the~~The present invention ~~is that~~includes a mold apparatus for powder molding, ~~comprising~~having a mold body with a through-hole for forming a side of a powder molding product, a lower punch to be fitted into the through-hole from beneath, an upper punch to be fitted into the through-hole from above, a spray member from which a lubricant solution is faced into the through-hole, a heater provided around a molding portion of the powder molding product, the molding portion being defined by the through-hole and the lower punch which is fitted into the through-hole, a temperature control system keeping a temperature of the heater higher than an evaporating temperature of the aqueous solution, and if required lower than a melting temperature of said lubricant; and an aqueous solution in which water soluble lubricant having at least 3g of solubility for 100 g of water at 20°C is dissolved in water, is provided in said spray member. By forming a crystallized layer of lubricant on a molding portion, the reduction of a pressure for ejecting the powder molding product, or the improvement of the density of the powder molding product can be obtained. Further, consecutive molding can be stably carried out.

Fig. 1 is a cross sectional diagram showing a first process according to an Example 1 of the present invention.

Fig. 2 is a cross sectional diagram showing a second process according to an Example 1 of the present invention.

5 Fig. 3 is a cross sectional diagram showing a third process according to an Example 1 of the present invention.

Fig. 4 is a cross sectional diagram showing a forth process according to an Example 1 of the present invention.

Fig. 5 is a solubility diagram of a soap.

10 Fig. 6 is a cross sectional diagram showing a first process according to an Example 2 of the present invention.

Fig. 7 is a cross sectional diagram showing a second process according to an Example 2 of the present invention.

15 Fig. 8 is a cross sectional diagram showing a third process according to an Example 2 of the present invention.

Fig. 9 is a cross sectional diagram showing a fourth process according to an Example 2 of the present invention.

Fig. 10 is a cross sectional diagram showing a first process according to an Example 3 of the present invention.

20 Fig. 11 is a cross sectional diagram showing a second process according to an Example 3 of the present invention.

Fig. 12 is a cross sectional diagram showing a first process according to an Example 4 of the present invention.

25 Fig. 13 is a cross sectional diagram showing a second process according to an Example 4 of the present invention.

~~Best Mode for Carrying Out~~Detailed Description of the In-
vention

Suitable embodiments of the present invention will
5 now be explained with reference to attached drawings. Further, it should be understood that the embodiments explained below are not limited to the scope of the present invention described in claims. And, it should be understood that the total constitution described below is not
10 essential requirement for the present invention.

Example 1

Example 1 of the present invention will now be explained with reference to Figs. 1 to 4. Fig. 1 represents
15 a first process. According to the same Figures, numeral 1 designates a through-hole formed in a die 2 serving as a mold for molding sides of a powder molding product A, i.e., compact as a later-described powder molded body. A lower punch 3 is fitted into the through-hole from the underneath
20 thereof and an upper punch 4 is also fitted into the through-hole ~~2-1~~ from the above thereof. A feeder 5, which provides a raw powder M, is slidably provided on an upper surface of the die 2. Above the through-hole ~~2-1~~ is provided a spray member 6 serving as a solution applying means
25 for spraying a lubricant solution L so as to attach the

same to a molding portion 1A of the mold. The spray member 6 is arranged so as to face the through-hole ~~2~~1, and is connected to a tank of the solution L (not shown) via an automatically openable and closable valve (not shown). A
5 heater 7 and a temperature detector 8 are provided around the periphery of the molding portion 1A for forming the powder molding product A, the molding portion being defined by the through-hole 1 and the lower punch 3 which is fitted into the through-hole. The heater 7 and the temperature de-
10 tector 8 are connected to a temperature control device 9 serving as a temperature controlling means. By the temperature control device 9, the temperature of through-hole ~~2~~1 is kept higher than the evaporating temperature of the aqueous solution L, and lower than the melting temperature
15 of the lubricant.

In the first process, due to the heat of the heater 7 being pre-controlled by the temperature control system 9, the temperature of the periphery of the through-hole 1 is kept higher than the evaporating temperature of the aqueous
20 solution L, and lower than the melting temperature of the lubricant. Then, the automatically openable and closable valve is opened to apply the aqueous solution L in which water soluble lubricant having at least 3g of solubility for 100 g of water at 20°C is dissolved in water by spray-
25 ing from the spray member 6 to the molding portion 1A of

the die 2 heated by the heater 7, with the lower punch 3 being fitted into the through-hole 1 to define the molding portion 1A. As a result, the solution L is evaporated and dried out, and thus crystals are allowed to grow on the peripheral surface of the through-hole 1, so that a crystallized layer B of the lubricant is uniformly formed. Using a water soluble lubricant having at least 3g of solubility for 100 g of water at 20°C can be produced precipitate in an aqueous solution around the room temperature. Therefore, trouble such as the clogging of the spray member 6 is occurred when applying by spray member 6. Meanwhile, for example, if an aqueous solution does not ~~keeping~~keep a higher temperature, conventional higher fatty acidic soaps comprising sodium stearate, sodium palmitate, sodium myristate, sodium laurate cannot be obtained at least 3g of solubility for 100 g of water.

Next, as illustrated in a second process shown in Fig. 2, the feeder 5 is moved forward so as to drop a raw powder M into the molding portion 1A to fill the same therewith. Subsequently, as illustrated in a third process shown in Fig. 3, the die 2 is moved downwardly, while the upper punch 4 is inserted into the molding portion 1A of the through-hole 1 from there above, so that the raw powder M is compressed in a manner that is sandwiched between the upper punch 4 and the lower punch 3. At this stage, a bot-

tom end of the lower punch 3 is firmly held in position. And in this third process, the material powder M is compressed as the state of lubrication by being pressed against the crystallized layer B formed of the lubricant

5 The powder molding product A thus press-molded becomes ejectable when the die 2 is moved further downwardly until the upper surface of the die 2 becomes essentially as high as the lower surface of the lower punch 3, as illustrated in a fourth process shown in Fig. 4. When ejecting
10 the same, the powder molding product A is allowed to contact the crystallized layer B that is formed of the lubricant and is in a lubricated condition. After ejecting the powder molding product A thus way, the first process is repeated and thus the aqueous solution L is applied to the
15 molding portion 1A again to form the crystallized layer B, and then the raw powder M is filled into the molding portion 1A.

Next, as to a good water solubility, it will be explained the point that the solubility is at least 3g of
20 solubility for 100 g of water at 20°C. As can be seen from the solubility for various fatty acid soaps illustrated in Fig. 4, the solubility of the mixed soaps which is produced by animal oil or vegetable oil or main components thereof are very lower at room temperature, thus even though it is
25 dissolved in water the precipitates are generated in a few

minutes. And at about 20 °C, which is used commonly as room temperature, the precipitates are generated. Therefore inconvenience such as the clogging of the spray member is occurred. In this regard, the recognition that these
5 component should not be included makes the solubility in 100 g water at 20°C is at least 3 g.

Preferred examples and comparative examples will now be explained with reference to Tables 1 to 3. In each of the preferred examples and comparative examples shown in
10 Tables 1 to 3, iron powders (average particle diameter: 90 μm) were used as the raw powder, to which was added 0.2% by weight of Lithium stearate (average particle diameter: 5 μm) serving as the lubricant, which were then mixed for 30 minutes using a rotary mixer, so that 7g of the resultant mixture of the raw powder was filled into a mold forming a cylindrical column having a 1 cm² pressurization area, and then 100 powder molding products were successively formed at a molding pressure of 8 t/cm². And in the examples, after the aqueous solution of the water-soluble lubricant
15 dissolved in water was applied to the molding portion heated at 150°C in the mold, it was evaporated and dried to form the crystallized layer, and then the raw powders were filled into this molding portion. In the comparative example 1, after the solution of lithium stearate (average particle diameter: 5 μm) dispersed in acetone was applied to
25

the molding portion of the mold heated at 150°C, it was dried to form the film, and then the raw powders were filled into this molding portion. The comparative example 2 is a case in which the lubricant was not applied to the
5 mold. Density R in each Table shows difference between maximum and minimum values in the density of 100 molding bodies continuously molded.

[Table 1]

	Example1	Example2	Example3	Example4	Example5	Example6	Example7	Example8	Example9
Mold lubricating component	Dipotassium hydrogen phosphate	Disodium hydrogen phosphate	Trisodium phosphate	Sodium polyphosphate	Riboflavin sodium phosphate	potassium sulfate	sodium sulfite	sodium thiosulfate	Sodium dodecylsulfate
Solvent	Water	Water	Water	Water	Water	Water	Water	Water	Water
State of lubricating component	dissolved	dissolved	dissolved	dissolved	dissolved	dissolved	dissolved	dissolved	dissolved
Concentration	1%	1%	1%	1%	1%	1%	1%	1%	1%
Molding temperature	150°C	150°C	150°C	150°C	150°C	150°C	150°C	150°C	150°C
Average ejecting pressure	6kN	8 kN	6 kN	8 kN	20 kN	18kN	20 kN	18 kN	16 kN
Average molding product density	7.56 g/cm ³	7.55 g/cm ³	7.56 g/cm ³	7.54 g/cm ³	7.5 g/cm ³	7.52 g/cm ³	7.5 g/cm ³	7.51 g/cm ³	7.53 g/cm ³
Density R	0.02	0.02	0.02	0.02	0.03	0.02	0.02	0.02	0.03

[Table 2]

	Example10	Example11	Example12	Example13	Example14	Example15	Example16	Example17	Example18
Mold lubricating component	Sodium dodecylbenzen e-sulfonate	Food Blue No.1	Food Yellow No.5	Sodium ascorbyl sulfate	sodium tetraborate	sodium silicate	sodium tungstate	sodium acetate	Sodium benzoate
Solvent	Water	Water	Water	Water	Water	Water	Water	Water	Water
State of lubricating component	dissolved	dissolved	dissolved	dissolved	Dissolved	dissolved	dissolved	dissolved	dissolved
Concentration	1%	1%	1%	1%	1%	1%	1%	1%	1%
Molding temperature	150℃	150℃	150℃	150℃	150℃	150℃	150℃	150℃	150℃
Average ejecting pressure	16kN	16 kN	20 kN	8 kN	8 kN	10kN	12 kN	18 kN	10 kN
Average molding product density	7.53 g/cm ³	7.53 g/cm ³	7.51 g/cm ³	7.54 g/cm ³	7.54 g/cm ³	7.54 g/cm ³	7.53 g/cm ³	7.51 g/cm ³	7.54 g/cm ³
Density R	0.02	0.03	0.04	0.02	0.02	0.03	0.03	0.02	0.02

[Table 3]

	Example19	Example21	Example23	Example24	Example25	Comparative Example 1	Comparative Example 2
Mold lubricating component	Sodium terephthalate	Sodium stearate	Sodium hydrogen carbonate	Sodium carbonate	Potassium nitrate	Lithium stearate	None
Solvent	Water	Water	Water	Water	Water	acetone	
State of lubricating component	dissolved	dissolved	dissolved	dissolved	dissolved	dispersed	
Concentration	1%	0.2%	1%	1%	1%	1%	
Molding temperature	150°C	150°C	150°C	150°C	150°C	150°C	150°C
Average ejecting pressure	1kN	16 kN	18 kN	18 kN	20 kN	22kN	32 kN
Average molding product density	7.54 g/cm ³	7.52 g/cm ³	7.51 g/cm ³	7.52 g/cm ³	7.51 g/cm ³	7.5 g/cm ³	7.48 g/cm ³
Density R	0.02	0.04	0.03	0.02	0.04	0.02	0.16

Comparison result from Tables 1 to 3 indicates that the pressure required for ejecting a compact from a die in the examples were less than or equal to that of the comparative example 1. Besides, the densities were improved in the examples as compared to the comparative example 1. Moreover, the densities R in the examples noticeably became smaller than that of the comparative example 1. Therefore, it is apparent from the result that the molding can be stably carried out according to the examples, even though it is carried out successively.

As is clearly indicated in Tables 1 to 3, the afore-said lubricant may suitably be a water soluble phosphate based metal salt, and the one having a phosphate group in its structure, such as dipotassium hydrogen phosphate, disodium hydrogen phosphate, tripotassium phosphate, trisodium phosphate, potassium polyphosphate, sodium polyphosphate, riboflavin potassium phosphate, riboflavin sodium phosphate or the like.

As is also seen from Tables 1 to 3, it is suitable that, as a water soluble sulfate-based metal salt, the lubricant may include a sulfate-based group in its structure, such as potassium sulfate, sodium sulfate, potassium sulfite, sodium sulfite, potassium thiosulfate, sodium thiosulfate, potassium dodecyl sulfate, sodium dodecyl sulfate, potassium dodecylbenzenesulfate, sodium dodecylbenzenesul-

fate, Food Blue No.1. ($C_{37}H_{34}N_2Na_2O_9S_3$), Food Yellow No.5. ($C_{16}H_{10}N_2Na_2O_7S_2$), potassium ascorbyl sulfate, sodium ascorbyl sulfate.

As is also seen from Tables 1 to 3, it is suitable
5 that, as a water soluble borate based metal salt, the lubricant may include a borate-based group in its structure, such as potassium tetraborate, sodium tetraborate.

Tables 1 to 3 also show that as a water soluble silicate based metal salt, the lubricant may suitably include a
10 silicate-based group in its structure, such as potassium silicate, sodium silicate.

Still also, Tables 1 to 3 show that as a water soluble tungstate based metal salt, the lubricant may suitably include a tungstate-based group in its structure, such as
15 potassium tungstate or sodium tungstate.

Table 1 to 3 show that as a water soluble organic acid based metal salt, the lubricant may suitably include an organic acid based group in its structure, such as potassium acetate, sodium acetate, potassium benzoate, sodium
20 benzoate, dipotassium terephthalate, disodium terephthalate, potassium ascorbate, or sodium ascorbate.

It is also seen from Tables 1 to 3, that as a water soluble nitrate based metal salt, the lubricant may suitably include a nitrate-based group in its structure such as
25 potassium nitrate, sodium nitrate.

It is still also seen from Tables 1 to 3 that as a water soluble carbonate-based metal salt, the lubricant may suitably include a carbonate based group in its structure, such as potassium carbonate, sodium carbonate, potassium
5 hydrogen carbonate or sodium hydrogen carbonate.

Alternatively, one or more of the foregoing lubricants may be used as the lubricant.

And the water soluble lubricant should have a concentration greater than or equal to 0.01 % by weight but less
10 than a concentration of a saturation. This is because the concentration of less than 0.01 % by weight makes it difficult to obtain a stably forming with constant temperature and speed since water content to apply and evaporate on the mold for forming is too large quantities and thus the mold
15 temperature is lower, while at the saturated concentration or above it does not allow the lubricant to be completely dissolved so that it is precipitated as a solid, thus causing troubles such as the clogging of the spray member 6 when applying lubricant by a spray member 6.

20 For dissolvent water, water from which metal and halogen elements are removed is preferable, such as distilled water or ion exchange water. This is because some lubricants, though it depends on a kind thereof, are precipitated due to the readiness to substitute metal components in water, thus causing troubles, while water contain-
25

ing a large amount of halogen components is likely to cause a bond to a compact or to produce a harmful substance such as dioxin or the like during a sintering process.

Further, some lubricants, though also depending on a
5 kind thereof, facilitate the growing of microorganisms and thus the solution is easily decayed, thereby causing a change in components, emitting bad smell. However, adding an antiseptic agent can prevent the growing of microorganisms. For the antiseptic agent, it is preferable to use one
10 which does not impair lubrication property, produces low harmful effects to a human body, and includes no halogen components, such as sodium benzoate or the like.

Furthermore, some lubricants have a problem that foaming easily occurs, and thus when the aqueous solution
15 (L) is applied to the molding portion (1A), such molding is likely to occur foaming so that a raw powder is caked. However, by adding a water-soluble solvent such as alcohol or ketone, or a defoaming agent, such foaming can be prevented. For alcohol or ketone, it is preferable to use one which
20 does not impair the lubricating action, causes less damages to a human body, and does not include halogen components, such as ethanol, acetone or the like.

In some cases, using a water soluble solvent such as alcohol and ketone with a lower boiling point or a lower
25 latent heat of evaporation than water can reduce hours for

evaporation or dry, eliminating the need for keeping the mold body 2 at high temperature.

In a case where these lubricants, additives or dis-
solvent water include halogen elements, a substance that is
5 highly toxic even in minute amounts such as dioxin is
likely to be created under such a condition that sintering
is performed with carbon components being coexistent, as is
often used in powder metallurgy of iron. Therefore it is
preferable to include no halogen elements therein.

10 As for the temperature of the mold body 2 and the
mixed raw powder M, keeping them at high temperature is de-
sirable because it contributes to reduction of hours for
drying, accompanied by effects of warm molding and the like.
If there is caused no particular trouble, however, it can
15 be kept at ordinary temperature. On the other hand, when
setting them at high temperature, it is preferable to
choose such a lubricant that is not melt down at a preset
temperature, since the melt lubricant makes it difficult to
stably perform warm molding due to the melt lubricant cak-
20 ing a raw powder, flowing down to the bottom of the mold
(the molding portion 1A). If there is caused no particular
trouble, however, it may be in a semi-molten state, in a
highly viscous state, or otherwise, at least one lubricant
of the mixed two ore more lubricants may be in a molten
25 state. Since zinc stearate and lithium stearate that have

been conventionally used have melting temperatures of about 120°C and about 220°C, respectively, it has heretofore been difficult to stably perform warm molding at a temperature higher than these temperatures. Among the lubricants proposed in the present invention, however, there are a number of lubricants that have a higher melting point than 220 °C, and some of them have a higher melting point than 1000°C. Therefore it is possible to easily and stably perform warm molding by raising the temperature almost to an oxidization temperature of the raw powder or heat resistance of mold (molding portion A). In that case, however, there occur problems such as fluidity of the raw powder, and thus it is preferable to use the lubricant that does not melt even under high temperature, as the one to be added into the mixed raw powder M. For example, the powdery lubricants of the present invention or solid lubricants such as graphite or molybdenum disulfide which can be used a high temperature of at least 200°C are preferable. Alternatively, it is also preferable to mold only by lubrication of the mold itself without using the lubricant into raw powder.

According to the description of the foregoing embodiment, there is provided a method for forming a powder molding product, comprising a filling the molding portion 1A in the mold body 2 with the raw powder M, and then inserting upper and lower punches 3, 4 into the molding portion 1A to

thereby form the powder molding product, wherein prior to filling the molding portion 1A with the raw powder M, the aqueous solution L in which lubricant is dissolved in a solvent is applied to the molding portion 1A to a uniform phase, and then the aqueous solution L is evaporated to thereby form a crystallized layer B on the molding portion 1A. Thus, the fine and uniform layer B for lubrication is formed on the peripheral surface of the molding portion 1A, thereby enabling the reducing of a pressure required for ejecting the powder molding product A from the molding portion 1A as well as the improving of the density of the powder molding product A.

Also, there is provided a mold apparatus for powder molding, comprising the mold body 2 with the through-hole 1 for molding a side of the powder molding product A, the lower punch 3 to be fitted into the through-hole 1 from beneath, the upper punch 4 to be fitted into the through-hole 1 from above, the spray member 6 from which the lubricant aqueous solution L is faced into the through-hole 1, the heater 7 provided around the molding portion 1A of the powder molding product A, the molding portion 1A being defined by the through-hole 1 and lower punch 3 which is fitted into the through-hole 1, and the temperature control system 9 keeping a temperature of the heater 7 higher than an evaporating temperature of the aqueous solution L, if re-

quired but lower than a melting temperature of said lubricant. And prior to filling the molding portion 1A with the raw powder M, the lubricant aqueous solution L is applied to the molding portion 1A which is heated, and then the aqueous solution L is evaporated to thereby form the fine and uniform crystallized layer B on the peripheral surface of the molding portion 1A. Thus, the fine and uniform crystallized layer B for lubrication is formed on the peripheral surface of the molding portion 1A, thereby enabling the reducing of a ~~pressure~~pressure required for ejecting the powder molding product A from the molding portion 1A as well as the improving of the density of the powder molding product A, and realizing the stable and successive molding.

Example 2

Fig.5-Fig.8 represent example 2. in which the same reference symbols as Example 1 will be designated by the same symbols, and their repeated detailed description will be omitted. A surface 10 of the through-hole 1 is formed with a surface treatment layer 11 by hydrophilicity imparting treatment to the surface 10 for improving the wetting action of the aqueous solution L relative to the surface 10, or by arranging hydrophilic material thereon. An angle X of contact of the surface treatment layer 11 relative to the aqueous solution L is smaller than an angle Y of contact of

the surface 10, which is made from the material of the die 2 itself, or of the upper surface 2A where the material is exposed, relative to the aqueous solution L ($X < Y$), thus enabling the said wetting action to be improved. Further, it should be noted herein that these angles of contact X, Y are not measured under such condition as shown in figure which are only schematically illustrated for the sake of explanation, but are measured under an equal condition, such as keeping the surface 10 and the upper surface 2A horizontally. And the surface treatment layer 11 is formed by: the thermal spraying, PVD, CVD or shot peening of oxide, fluoride, nitride, chloride, sulfide, bromide, iodide, carbide, hydroxide and etc. having bonds as shown in Table 4 to hydrophilic coating, subjecting the coating of titania, zinc oxides or the like to photocatalytic reaction by irradiating light thereto, creating hydroxide by alkali or hydrothermal treatment, the surface treatment by sputtering with potassium ions or sodium ions, and utilizing change in surface tension of the aqueous solution L by the formation of minute pores on the surface by spray coating or powder metallurgy die, whereby the surface treatment layer thus obtained allows the angle of contact of the solution relative to the surface 10 of the through-hole 1 to be made smaller, thereby improving the wetting action of the solution therein. Alternatively, the surface 10 of the through-

hole 1 may undergo the removal of oily organisms through acid or flame processing, electrolytic polishing etc so that the angle of contact α may become small. If there causes no problem in strength, the die may preferably be
5 formed from hydrophilic materials shown in Table 4. Alternatively, metals such as iron or hard metal may have the substances shown in Table 1 dispersed therein to improve strength and hardness. Alloying with easily oxidizable metals such as Ti, V, Si, and Al, etc. to use as the material
10 of the die is also effective to improve hydrophilic property. In the case of coating, the coating of iron or hard metal together with hydrophilic materials in order to improve strength and hardness is desirable since such coating can satisfy both the long-duration and hydrophilicity of
15 the die.

[Table 4]

Examples of Hydrophillic Substances				
Hydrophillic Bond Elements or Hydrophillic Substances			Approximate ionicity Bond	Principal Reason for Hydrophillic Property
Cs-F, Fr-F			93%	due to large ionicity (polarity) of Bonds
K-F, Rb-F			92%	
Na-F, Ba-F, Ra-F			91%	
Li-F, Ca-F, Sr-F			89%	
Ac-F, lanthanoid-F			88%	
Mg-F, Y-F, Cs-O, Fr-O			86%	
Se-F, Hf-F, Th-F, K-O, Rb-O			84%	
Zr-F, Pa-F, U-F, Na-O, Ba-O, Ra-O			82%	
Be-F, Al-F, Ti-F, Ta-F, Mn-F, Li-O, Ca-O, Sr-O			79%	
Nb-F, V-F, Cr-F, Zn-F, Ga-F, Ac-O, lanthanoid-O			76%	
W-F, Cd-F, In-F, Mg-O, Y-O, Cs-O, Fr-O, Cs-N, Fr-N, Cs-Cl, Fr-Cl			73%	
Mo-F, Fe-F, Ti-F, Si-F, Ge-F, Sn-F, Se-O, Hf-O, Th-O, K-N, Rb-N, K-Cl, Rb-Cl			70%	
Re-F, Tc-F, Co-F, Ni-F, Cu-F, Ag-F, Hg-F, Pb-F, Bi-F, Zr-O, Pa-O, U-O, Na-N, Ba-N, Ra-N, Na-Cl, Ba-Cl, Ra-Cl, Cs-Br, Fr-Br			67%	
B-F, As-F, Po-F, Be-O, Al-O, Ti-O, Ta-O, Mn-O, Li-N, Ca-N, Sr-N, Li-Cl, Ca-Cl, Sr-Cl, K-Br, Rb-Br			63%	
P-F, Te-F, Nb-O, V-O, Cr-O, Zn-O, Ga-O, Ac-N, lanthanoid-N, Ac-Cl, lanthanoid-Cl, Na-Br, Ba-Br, Ra-Br				
P-F, Te-F, Nb-O, V-O, Cr-O, Zn-O, Ga-O, Ac-N, lanthanoid-N, Ac-Cl, lanthanoid-Cl, Na-Br, Ba-Br, Ra-Br			59%	
Ru-F, Os-F, Rh-F, Ir-F, Pt-F, Au-F, W-O, Cd-O, In-O, Mg-N, Y-N, Cs-N, Fr-N, Mg-Cl, Y-Cl, Cs-Cl, Fr-Cl, Li-Br, Ga-Br, Sr-Br, Cs-C, Fr-C, Cs-S, Fr-S, Cs-I, Fr-I			55%	
Mo-O, Fe-O, Ti-O, Si-O, Ga-O, Sn-O, Se-N, Hf-N, Th-N, Se-Cl, Hf-Cl, Th-Cl, Ac-Br, lanthanoid-Br, K-C, Rb-C, K-S, Rb-S, K-I, Rb-I			51%	
Au-F, Se-F, Re-O, Tc-O, Co-O, Ni-O, Cu-O, Ag-O, Hg-O, Pb-O, Sb-O, Bi-O, Zr-N, Pa-N, U-N, Zr-Cl, Pa-Cl, U-Cl, Mg-Br, Y-Br, Na-C, Ba-C, Ra-C, Na-S, Ba-S, Ra-S, Na-I, Ba-I, Ra-I			47%	
B-O, As-O, Po-O, Be-N, Al-N, Ti-N, Ta-N, Mn-N, Be-Cl, Al-Cl, Ti-Cl, Ta-Cl, Mn-Cl, Se-Br, Hf-Br, Th-Br, Li-C, Ga-C, Sr-C, Li-S, Ca-S, Sr-S, Li-I, Ga-I, Sr-I			43%	
P-O, Ta-O, Nb-N, V-N, Cr-N, Zn-N, Ga-N, Nb-Cl, V-Cl, Cr-Cl, Zn-Cl, Ga-Cl, Zr-Br, Pa-Br, U-Br, Ac-C, lanthanoid-C, Ac-S, lanthanoid-S, Ac-I, lanthanoid			39%	
Ru-O, Os-O, Rh-O, Ir-O, Pd-O, Pt-O, Al-O, W-N, Cd-N, In-N, W-Cl, Cd-Cl, In-Cl, Be-Br, Al-Br, Ti-Br, Ta-Br, Mn-Br, Mg-C, Y-C, Cs-C, Fr-C, Mg-S, Y-S, Cs-S, Fr-S, Mg-I, Y-I, Cs-I, Fr-I			35%	
Mo-N, Fe-N, Ti-N, Si-N, Ge-N, Sn-N, Mo-Cl, Fe-Cl, Ti-Cl, Si-Cl, Ge-Cl, Sn-Cl, Nb-Br, V-Br, Cr-Br, Zn-Br, Ga-Br, Se-C, Hf-C, Th-C, Se-S, Hf-S, Th-S, Se-I, Hf-I, Th-I			30%	
General Substances including Hydroxyl Group				due to the inclusion of hydroxyl group
Oxides in General				due to surface being termed to include hydroxyl
Water-Soluble Substances in General				due to being soluble in water
Some Specific Oxides (e.g., Titanium oxide, zinc oxide)				due to photo-excitation

And in the first process, due to the heat of the heater 7 being pre-controlled by the temperature control system 9, the temperature of the surface 10 of the through-hole 1 is kept higher than the evaporating temperature of the aqueous solution L, and lower than the melting temperature of the lubricant beforehand. Then, the automatically openable and closable valve is opened to apply the aqueous solution L of the lubricant by spraying from the spray member 6 to the molding portion 1A of the die 2 heated by the heater 7, with the lower punch 3 being fitted into the through-hole 1 to define the molding portion 1A. At this moment, the angle X of contact of the aqueous solution L, which would be the angle Y of contact without the surface treatment layer 11, is allowed to be the smaller angle X of contact owing to the surface treatment layer 11, thus allowing the aqueous solution L to be prevented from being repelled, to thereby the aqueous solution L is applied to the entire surface of the through-hole 1 and wet the same. As a result, the aqueous solution L is evaporated and dried out, and thus crystals are allowed to grow entirely on the surface treatment layer 11 of the through-hole 1, so that a crystallized layer B serving as a lubricating layer of the lubricant is uniformly formed.

Next, as illustrated in a second process shown in Fig. 6, the feeder 5 is moved forward so as to drop a raw powder

M into the molding portion 1A to fill the same therewith. Subsequently, as illustrated in a third process shown in Fig. 7, the die 2 is moved downwardly, while the upper punch 4 is inserted into the molding portion 1A of the through-hole 1 from there above, so that the raw powder M is compressed in a manner that is sandwiched between the upper punch 4 and the lower punch 3. At this stage, a bottom end of the lower punch 3 is firmly held in position. In this third process, the raw powder M is compressed by being pressed against the crystallized layer B which is formed by lubricant in a lubricated condition.

The powder molding product A thus press-molded becomes ejectable when the die 2 is moved further downwardly until the upper surface of the die 2 becomes essentially as high as the upper surface of the lower punch 3, as illustrated in a fourth process shown in Fig. 9. When ejecting the same, the powder molding product A is allowed to contact the crystallized layer ~~1-E~~E which is formed by lubricant in a lubricated condition. After ejecting the powder molding product A thus way, the first process is repeated and thus the aqueous solution L is applied to the molding portion 1A again to form the crystallized layer ~~1-E~~E, and then the raw powder M is filled into the molding portion 1A.

As is apparent from the foregoing, the surface 10 of the through-hole 1 is formed with the surface treatment

layer 11 so as to have the smaller angle X of contact with the aqueous solution L than the angle Y of contact of the die 2 with the aqueous solution L, in accordance with the foregoing experiment. Thus, when the aqueous solution L is applied, the wetting action of the aqueous solution L relative to the ~~through-hole~~surface 10 is improved so that the aqueous solution L can be extended over the surface treatment layer 11, eventually over the entire surface of the through-hole. Consequently, the entire surface of through-hole 1 can be formed with the crystallized layer B by performing water evaporation. As a result, high-density powder molding product A can be stably obtained.

Example 3

Fig. 9 and Fig. 10 represent Example 3, in which the same reference symbols as those in Example 1 and Example 2 will be designated by the same symbols, and their repeated detailed description will be omitted. According to the Example 2, the upper surface 2A of the die 2 on which feed 5 is slidably provided is formed with a surface treatment layer 21 by water repellency imparting treatment to the surface 2A for improving its liquid repelling ability (i.e., reducing the wetting action of the aqueous solution L) relative to the surface 2A, or by arranging water repellent material thereon. An angle Y' of contact of the surface

treatment layer 21 relative to the aqueous solution L is larger than an angle X' of contact of the surface made from the material of the die 2 itself, or in Example 3 the surface 10 of the through-hole 1, relative to the aqueous solution L ($Y' > X'$), thus enabling the said wetting action to be reduced. The above surface treatment layer 21 may be formed from silicone- or fluorine-based resin such as those including Si-H bond, or C-H bond, etc., or from nonpolar substances, as shown in Table 5.

10

[Table 5]

Examples of Water Repellent Substances

Water Repellent Bond elements or Water Repellent Substances	Approximate ionicity of Bond	Principal Reason for Water Repellency
Re-H, Ic-H, Co-H, Ni-H, Cu-H, Ag-H, Hg-H	1%	due to small ionicity (polarity) of Bonds
Mo-H, Fe-H, Ti-H, Si-H	3%	
H-C, P-C, Ie-C, H-S, P-S, Ie-S, H-I, P-I, Ie-I, W-H, Cd-H, In-H	4%	
B-C, As-C, Po-C, B-S, As-S, Po-S, B-I, As-I, Po-I, Nb-H, V-H, Cr-H, Zn-H, Ga-H	7%	
Re-C, Ic-C, Co-C, Ni-C, Cu-C, Ag-C, Hg-C, Pb-C, Sb-C, Bi-C, Re-S, Ic-S, Co-S, Ni-S, Cu-S, Ag-S, Hg-S, Pb-S, Sb-S, Bi-S, Re-I, Ic-I, Co-I, Ni-I, Cu-I, Ag-I, Hg-I, Pb-I, Sb-I, Bi-I, Be-H, Al-H, Li-H, Ia-H, Mn-H	9%	
Mo-C, Fe-C, Ti-C, Si-C, Ge-C, Sn-C, Mo-S, Fe-S, Ti-S, Si-S, Ge-S, Sn-S, Mo-I, Fe-I, Ti-I, Si-I, Ge-I, Sn-I, Zr-H, Pa-H, U-H	11%	due to being nonpolar
Nonpolar Substances in General		

According to Example 3, therefore, the automatic open
able and closable valve is opened so that the aqueous solu-
tion L of the lubricant is sprayed from the spray member 6
and applied to the molding portion 1A of the die 2 that is
5 heated by the heater 7. At this moment, part of the aqueous
solution L is likely to be attached to the upper surface 2A
of the die 2. Nevertheless, the aforementioned angle Y' of
contact of the aqueous solution L with the upper surface 2A
on which the surface treatment layer 21 is provided, be-
10 comes larger than the angle X' of direct contact thereof
with the die 2, whereby the aqueous solution L is allowed
to be repelled, thus preventing the aqueous solution L to
collect on the surface 2A.

As is apparent from the foregoing, since the upper
15 surface 2A is formed with the surface treatment layer 21 so
as to have the larger angle Y' of contact with the aqueous
solution L than the angle X' of contact of the die 2 itself
with the aqueous solution L, whereby the water repellent
property on the upper surface 2A can be improved, making
20 the aqueous solution L less likely to pile up or collect on
the upper surface 2A (the surface treatment layer 21), thus
preventing the aqueous solution L from collecting on the
upper surface 2A (surface treatment layer 21), which in
turn makes the raw powder M housed in the feeder 5 less

likely to be contacted by the aqueous solution L, thereby enabling the raw powder M to be prevented from caking.

Example 4

Fig. 9 and Fig. 10 represent Example 4, in which the same reference symbols as those in Examples 1~3 will be designated by the same symbols, and their repeated detailed description will be omitted.

According to example 4, above the through-hole 1 is provided the spray member 6 serving as applying means for spraying the aqueous solution L so as to attach the aqueous solution L to the molding portion 1A. The spray member 6 is arranged so as to face the through-hole 1. The aqueous solution L contains components which improve the wetting action relative to the surface 10 of the through-hole 1. The wetting action improving components are ones that can make the angle X'' of contact of the aqueous solution L with the surface 10 smaller, for example, surfactant is used.

Thus, the automatically open able and closable valve is opened to apply the aqueous solution L of the lubricant by spraying from the spray member 6 to the molding portion 1A of the die 2 heated by the heater 7, with the lower punch 3 being fitted into the through-hole 1 to define the molding portion 1A. At this moment, the angle X'' of contact of the aqueous solution L, which would become large without the wetting action improving components, is allowed to be

small enough owing to the components, thus allowing the aqueous solution L to be prevented from being repelled, to thereby be applied to the entire surface 10 of the through-hole 1 and wet the same. As a result, the aqueous solution
5 L is evaporated and dried out, and thus crystals are allowed to grow entirely around the surface of the through-hole 1, so that a crystallized layer B of the lubricant is uniformly formed.

As is apparent from the foregoing, since the aqueous
10 solution L contains components which improve the wetting action in order to decrease the angle X'' of contact with the surface 10, the wetting action of the aqueous solution L in the through-hole 1 is improved when the aqueous solution L is applied, thus allowing the aqueous solution L to
15 be extended over the entire surface of the through-hole 1, so that the aqueous solution L is evaporated to thereby allow the crystallized layer B to grow entirely, thus enabling the high-density powder molding product to be stably obtained.

20 Further, hereinafter examples and comparative examples will now be explained with reference to Table 6. In each of the examples and comparative examples shown in Table 6, iron powders (average particle diameter: 90 μm) were used as the raw powder, and 7 g of the mixture of the raw
25 powder was filled into a mold forming a cylindrical column

having a 1 cm² pressurization area, and then powder molding products were formed at a molding pressure of 8 t/cm². And in the preferred examples, 1% solution of dipotassium hydrogen phosphate as water-soluble lubricant was applied to the molding portion of the die coated with hydrophilic material and heated to 250 °C, and then it was evaporated and dried out to form the crystallized layer, and then the raw powders were filled into this molding portion. In the comparative example 1, after the lubricant was applied to the molding portion of an ordinary die heated to 250 °C, it was dried and then the raw powder was filled into this molding portion. In the comparative example 2, after the lubricant was applied to the molding portion of an ordinary die heated to 150 °C, it was dried and then the raw powder was filled into the molding portion. The comparative example 3 is a case in which an ordinary die was heated to 150 °C, and then the raw powder was filled into the molding portion without the application of lubricant. In either example, SKH-51 as typically employed for tool steel was used for the molding portion of such ordinary die.

[Table 6]

	Example1	Example2	Example3	Example4	Example5	Example 6	Comparative Example 1	Comparative Example 2	Comparative Example 3
Hydrophilic Bond Element	Al-O Ti-O	Al-O	Ti-O	Al-O Mg-O	Al-O Si-O	Al-O Ca-O	None	None	None
Components of Hydrophilic Coating	Al ₂ O ₃ 60 % TiO ₂ 40 %	Al ₂ O ₃	TiO ₂	Spinel	Al ₂ O ₃ 60 % SiO ₂ 40 %	Al ₂ O ₃ 60 % CaO 40 %	None	None	None
Process for treatment of Hydrophilic Coating	Spray coating	Spray coating	Spray coating	Spray coating	Spray coating	Spray coating	None	None	None
Lubrication of Mold	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	No
Molding Temperature	250 °C	250 °C	250 °C	250 °C	250 °C	250 °C	250 °C	150 °C	150 °C
Molding Density	7.68 g/cm ³	7.67 g/cm ³	7.68 g/cm ³	7.67 g/cm ³	7.68 g/cm ³	7.67 g/cm ³	unformable	7.58 g/cm ³	Unformable

Comparison result from Table 6 indicates that molding was found impossible if it was performed at 250 °C using dies without the hydrophilic coating, due to the lubricant being not fully attached to the molding portion. According to the Examples 1-6 where molding was performed, using dies with the hydrophilic coating, molding was found possible at temperature higher than 150°C, and it was found that high-density molding product denser than those formed at 150°C can be obtained.

10

~~Industrial Applicability~~

Further, the present invention is not limited to the foregoing embodiment but may be variously modified within the scope of the invention. In the foregoing embodiment, said aqueous solution is applied to the molding portion and then evaporated the aqueous solution to form the crystallized layer on the molding portion prior to filling the raw powder, and then the punches fitted into the molding portion to thereby forming the powder molding product, however, it is not always necessary to form the crystallized layer on the molding portion by applying the solution thereto and then evaporating the same, prior to filling the raw powder. For example, after forming a first powder molding product, a second powder molding product may be formed by filling a second raw powder, utilizing the crystallized layer formed

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when the first powder molding product is formed, without applying the aqueous solution to the molding portion, and then the aqueous solution may be applied to the molding portion prior to filling a third raw powder, and then the
5 aqueous solution is evaporated, to thereby form a second crystallized layer on the molding portion. The aqueous solution may be applied to the molding portion in such an intermittent continuance.